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(54) Title: FUNCTIONAL BACTERIAL/MAMMALIAN CYTOCHROME P450 CHIMERA

(57) Abstract

The present invention is directed to a chimeric DNA molecule which includes a first DNA molecule encoding a portion of a full length bacterial P450 protein and a second DNA molecule fused to the first DNA molecule and encoding a portion of a full length mammalian P450 protein. The chimeric DNA molecule encodes a fusion protein which is active and soluble in aqueous liquid. A further aspect of the present invention is directed to the fusion protein encoded by the chimeric DNA molecule. The fusion protein is useful in bioremediation processes and also can be used to hydroxylate a compound to be oxidized.

FUNCTIONAL BACTERIAL/MAMMALIAN CYTOCHROME P450 CHIMERA

The subject matter of this application was made with support from the
United States Government National Institutes of Health Grant No. GM624(PPG),
ES060062, and ES05407. The Government may have certain rights.

This application claims benefit of U.S. Provisional Patent Application Serial No. 60/056,754, filed August 20, 1997, which is hereby incorporated by reference.

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FIELD OF THE INVENTION

The present invention relates to a functional bacterial/mammalian cytochrome P450 chimera.

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BACKGROUND OF THE INVENTION

Cytochrome P450 ("P450") is a term used for a widely distributed group of unique heme proteins which form carbon monoxide complexes with a major absorption band at wavelengths around 450 nm. These proteins are enzymes which carry out oxidations involved in biosynthesis and catabolism of specific cell or body components, and in the metabolism of foreign substances entering organisms.

Oxygenating enzymes such as P450 appear to be fundamental cellular constituents in most forms of aerobic organisms. The activation of molecular oxygen and incorporation of one of its atoms into organic compounds by these enzymes are reactions of vital importance not only for biosynthesis, but also for metabolic activation or inactivation of foreign agents such as drugs, food preservatives and additives, insecticides, carcinogens and environmental pollutants.

In eukaryotic systems P450, and P450 dependent enzymes are known to act on such xenobiotics and pharmaceuticals as phenobarbitol, antipyrine, haloperidol and prednisone. Known substrates of environmental importance include compounds such as DDT, and a variety of polychlorinated biphenyls and polyaromatic hydrocarbons, as well as other halogenated compounds, including halobenzenes and chloroform.

P4502C9: Baculovirus-mediated Expression, Purification, Structural Characterization, Substrate Stereoselectivity, and Prochiral Selectivity of the Wild-Type and I359L Mutant Forms," Arch. Biochem. Biophys., 333:447-458 (1996); Waterman, M.S., "Heterologus Expression of Mammalian P450 Enzymes," Advances Enzymol., 68:37-66 (1994)) and peptitergents to improve solubility. (Sueyoshi et al., "Molecular Engineering of Microsomal P4502a-4 to a Stable, Water-Soluble Enzyme," Arch. Biochem. Biophys., 322:265-271 (1995)). In contrast, the crystal structures of a number of cytosolic bacterial P450s have been determined. These include P450_{cam}, P450_{bm3}, P450_{terp}, and P450_{eryF}. (Poulos et al., "The 2.6-Δ Crystal 10 Structure of Psudomonas putida Cytochrome P-450," J. Biol. Chem., 260:16122-16130 (1985); Poulos et al., "High-Resolution Crystal Structure P450cam," J. Mol. Biol., 195:685-700 (1987); Ravichandran et al., "Crystal Structure of Hemeprotein Domain of P450BM-3, a Prototype for Microsomal P450's," Science, 261:731-736 (1993); Hasemann et al., "Crystal Structure and Refinement of Cytochrome P450_{tem} at 15 2.3 Δ Resolution," J. Mol. Biol., 1169-1185 (1994); Haseman et al., "Structure and Function of Cytochrome P450: A Comparative Analysis of Three Crystal Structures," Structure, 3:41-62 (1995); Cupp-Vickery et al., "Preliminary Crystallographic Analysis of an Enzyme Involved in Erythromycin Biosynthesis: Cytochrome P450_{eryF}," Proteins, 20:197-201 (1994)). Since no detailed structural information has 20 been obtained for a mammalian P450 enzyme, all attempts to determine the effect of enzyme-substrate interactions have used the crystal structures from the soluble bacterial P450 enzymes. (Cupp-Vickery et al., "Preliminary Crystallographic Analysis of an Enzyme Involved in Erythromycin Biosynthesis: Cytochrome P450_{ervF}," Proteins, 20:197-201 (1994); Paulsen et al., Methods in Enzymology, 25 272:337-46 (1996)). While homology models can be constructed for the membrane-bound mammalian enzymes based on the bacterial enzymes, the very low sequence identities (<20%) mean that any resulting model is of low resolution. In fact, no information directly shows that mammalian and bacterial enzymes are structurally related.

The present invention is directed to overcoming the deficiencies of the prior art by forming a P450 protein which is soluble and active in aqueous liquid.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A is a model of the chimeric structure of the present invention. The blue region is from P450_{cam} and the red region is from CYP2C9. The chimera contains 3 substrate recognition sites from P450_{cam} and 3 from CYP2C9. Figure 1B shows the construction of a fused plasmid of P450_{cam} and CYP2C9.

Figure 2A is a CO-reduced differential spectrum of the fusion protein of the present invention. The preparation used corresponds to lane 2 in Figure 2B. Figure 2B shows an SDS-polyacrylamide gel electrophoresis of the chimera of the present invention expressed in *E. coli*. Lanes 1 and 2 show the fusion protein and lane 3 and 4 show P450_{cam} wild-type. Lane 1, 105,000g supernatant (3μg protein); lane 2, eluate from a hydroxylaapatite column (1.5 μg protein); lane 3, 105,000g supernatant (3 μg protein); lane 4, eluate from hydroxylapatite column (2.2 μg protein); lane 5, molecular marker. The gel was stained with Coomassie Brilliant Blue R250.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is directed to a chimeric DNA molecule which

20 includes a first DNA molecule encoding a portion of a full length bacterial P450

protein and a second DNA molecule fused to the first DNA molecule and encoding a

portion of a full length mammalian P450 protein. The chimeric DNA molecule

encodes a fusion protein which is active and soluble aqueous liquid. This chimeric

DNA molecule can have the nucleotide sequence corresponding to SEQ. ID. No. 1 as

follows:

atgacgactg aaaccataca aagcaacgcc aatcttgccc ctctgccacc ccatgtgcca 60

gagcacctgg tattcgactt cgacatgtac aatccgtcga atctgtctgc cggcgtgcag 120

gaggcctggg cagttctgca agaatcaaac gtaccggatc tggtgtggac tcgctgcaac 180

ggcggacact ggatcgccac tcgcggccaa ctgatccgtg aggcctatga agattaccgc 240

cacttttcca gcgagtgccc gttcatccct cgtgaagccg gcgaagccta cgacttcatt 300

cccacctcga tggatccgcc cgagcagcg cagtttcgtg cgctggccaa ccaagtggtt 360

ggcatgccgg tggtggataa gctggagaac cggatccagg agctggcctg ctcgctgatc 420

gagagcctgc gcccgcaagg acagtgcaac ttcaccgagg actacgccga acccttcccg 480

	Pr	o Gl	u Gl	n Ar 10	g Gli O	n Phe	e Ar	g Al	a Le	u Al 5	a As	n Gl:	n Vai	l Va 11		y Me
. 5	Pro	o Va	l Va 11	l As	p Lys	s Lei	ı Glı	120		g Il	e Gli	n Gli	u Let 125		a Cy	s Se
	Let	u Il 13	e Gl	u Se	r Leu	ı Arg	9 Pro		n Gl	y Gl	n Cys	3 Ası 140		ì Th:	r Glı	ı Asp
10	Ty:	r Al	a Gl	u Pro	o Phe	Pro 150	lle	e Arg	g Il	e Phe	8 Met		ı Lev	ı Ala	a Gly	/ Let
15	Pro	Gl	u Gli	u Asy	165	Pro	His	Lev	ı Ly:	170		ı Thi	Asp	Glr	1 Met	
	Arg	y Pro	o Asp	9 Gly 180	y Ser	Met	Thr	Phe	2 Ala 189		a Ala	Lys	Glu	190		туг
20	Asp	ту	r Let 195	ı Ile	Pro	Ile	Ile	200		n Arg	J Arg	Glr.	Lys 205		Gly	/ Asn
	Asn	210	Glr	n Asp	Phe	Ile	Asp 215		Phe	e Leu	Met	Lys 220		Glu	Lys	Glu
25	225				Pro	230					235					240
30	Ala	Va]	Asp	Leu	Phe 245	Gly	Ala	Gly	Thr	Glu 250		Thr	Ser	Thr	Thr 255	
				260					265					270		_
35			275		Ile			280					285			
		290			Ser		295					300				
40	305				Ile	310					315					320
45					Lys 325					330					335	
				340	Leu				345					350		
50	Asn	Pro	Glu 355	Met	Phe	Asp	Pro	His 360	His	Phe	Leu	Asp	Glu 365	Gly	Gly	Asn
	Phe	Lys 370	Lys	Ser	Lys		Phe 375	Met	Pro	Phe	Ser	Ala 380	Gly	Lys	Arg	Ile
55	Cys 385	Val	Gly	Glu	Ala	Leu . 390	Ala	Gly	Met	Glu	Leu 395	Phe	Leu	Phe	Leu	Thr 400
	Ser	Ile	Leu	Gln	Asn 405	Phe .	Asn	Leu	Lys	Ser 410	Leu	Val	Asp	Pro	Lys 415	Asn

Mammalian P450s on Basis of P450_{cam} X-ray Structure," Methods in Enzymology, 206:11-30 (1991), which is hereby incorporated by reference.

Suitable mammalian P450 proteins include 1A, 2B, 2C, 2D, and 3A families of cytochrome P450 and CYP2C9. CYP2C9, which is particularly preferred, has an amino acid sequence of SEQ. ID. No. 3 as follows:

Met Asp Ser Leu Val Val Leu Val Leu Cys Leu Ser Cys Leu Leu Leu 10 Leu Ser Leu Trp Arg Gln Ser Ser Gly Arg Gly Lys Leu Pro Pro Gly 25 Pro Thr Pro Leu Pro Val Ile Gly Asn Ile Leu Gln Ile Gly Ile Lys 15 Asp Ile Ser Lys Ser Leu Thr Asn Leu Ser Lys Val Tyr Gly Pro Val Phe Thr Leu Tyr Phe Gly Leu Lys Pro Ile Val Val Leu His Gly Tyr 20 Glu Ala Val Lys Glu Ala Leu Ile Asp Leu Gly Glu Glu Phe Ser Gly 25 Arg Gly Ile Phe Pro Leu Ala Glu Arg Ala Asn Arg Gly Phe Gly Ile Val Phe Ser Asn Gly Lys Lys Trp Lys Glu Ile Arg Arg Phe Ser Leu 30 Met Thr Leu Arg Asn Phe Gly Met Gly Lys Arg Ser Ile Glu Asp Arg Val Glu Glu Glu Ala Arg Cys Leu Val Glu Glu Leu Arg Lys Thr Lys 35 Ala Ser Pro Cys Asp Pro Thr Phe Ile Leu Gly Cys Ala Pro Cys Asn 40 Val Ile Cys Ser Ile Ile Phe His Lys Arg Phe Asp Tyr Lys Asp Gln Gln Phe Leu Asn Leu Met Glu Lys Leu Asn Glu Asn Ile Lys Ile Leu 195 45 Ser Ser Pro Trp Ile Gln Ile Cys Asn Asn Phe Ser Pro Ile Ile Asp Tyr Phe Pro Gly Thr His Asn Lys Leu Leu Lys Asn Val Ala Phe Met 50 225 230

Lys Ser Tyr Ile Leu Glu Lys Val Lys Glu His Gln Glu Ser Met Asp

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gcatggatat gaagcagtga aggaagccct gattgatctt ggagaggagt tttctggaag 300 aggcattttc ccactggctg aaagagctaa cagaggattt ggaattgttt tcagcaatgg 360 5 aaagaaatgg aaggagatcc ggcgtttctc cctcatgacg ctgcggaatt ttgggatggg 420 gaagaggagc attgaggacc gtgttcaaga ggaagcccgc tgccttgtgg aggagttgag 480 aaaaaccaag gcctcaccct gtgatcccac tttcatcctg ggctgtgctc cctgcaatgt 540 10 gatctgctcc attattttcc ataaacgttt tgattataaa gatcagcaat ttcttaactt 600 aatggaaaag ttgaatgaaa acatcaagat tttgagcagc ccctggatcc agatctgcaa 660 15 taatttttct cetateattg attactteec gggaacteac aacaaattac ttaaaaaegt 720 tgcttttatg aaaagttata ttttggaaaa agtaaaagaa caccaagaat caatggacat 780 gaacaaccct caggacttta ttgattgctt cctgatgaaa atggagaagg aaaagcacaa 840 20 ccaaccatct gaatttacta ttgaaagctt ggaaaacact gcagttgact tgtttggagc 900 tgggacagag acgacaagca caaccetgag atatgetete etteteetge tgaagcacce 960 25 agaggtcaca gctaaagtcc aggaagagat tgaacgtgtg attggcagaa accggagccc 1020 ctgcatgcaa gacaggagcc acatgcccta cacagatgct gtggtgcacg aggtccagag 1080 atacattgac cttctcccca ccagcctgcc ccatgcagtg acctgtgaca ttaaattcag 1140 30 aaactatete atteecaagg geacaaceat attaatttee etgaettetg tgetacatga 1200 caacaaagaa tttcccaacc cagagatgtt tgaccctcat cactttctgg atgaaggtgg 1260 35 caattttaag aaaagtaaat acttcatgcc tttctcagca ggaaaacgga tttgtgtggg 1320 agaagccctg geeggeatgg agetgttttt attectgace tecattttae agaactttaa 1380 cctgaaatct ctggttgacc caaagaacct tgacaccact ccagttgtca atggatttgc 1440 40 ctctgtgccg cccttctacc agctgtgctt cattcctgtc tgaagaagag cagatggcct 1500 ggctgctgct gtgcagtccc tgcagctctc tttcctctgg ggcattatcc atctttcact 1560 45 atctgtaatg cottttctca cotgtcatct cacattttcc ottccctgaa gatctagtga 1620 acattcgacc tocattacgg agagtttcct atgtttcact gtgcaaatat atctgctatt 1680 etecatacte tgtaacagtt geattgactg teacataatg eteatactta tetaatgttg 1740 50 agttattaat atgttattat taaatagaga aatatgattt gtgtattata attcaaaggc 1800 attictitic tgcatgttct aaataaaaag cattattatt tgctg 1845

Suitable bacterial P450 proteins include P450_{cam}, P450_{bm3}, P450_{terp}, and P450_{eryF}. These proteins are described in Poulos et al., "The 2.6-Δ Crystal Structure of *Psudomonas putida* Cytochrome P-450," J. Biol. Chem., 260:16122-16130 (1985); Poulos et al., "High-Resolution Crystal Structure P450cam," J. Mol.

60 Biol., 195:685-700 (1987); Ravichandran et al., "Crystal Structure of Hemeprotein

	Asp	210	a Ile	e Ser	: Ile	· Val	Ala 215	Asn	Gly	/ Gln	Val	Asn 220		Arg	Pro	Ile
5	Thr 225	Ser	Asp	Glu	Ala	Lys 230	Arg	Met	Cys	Gly	Leu 235		Leu	Val	Gly	Gly 240
	Leu	Asp	Thr	Val	Val 245	Asn	Phe	Leu	Ser	Phe 250		Met	Glu	Phe	Leu 255	
10	Lys	Ser	Pro	Glu 260	His	Arg	Gln	Glu	Leu 265	Ile	Glu	Arg	Pro	Glu 270		Ile
15	Pro	Ala	Ala 275	Cys	Glu	Glu	Leu	Leu 280	Arg	Arg	Phe	Ser	Leu 285	Val	Ala	Asp
	Gly	Arg 290	Ile	Leu	Thr	Ser	Asp 295	Tyr	Glu	Phe	His	Gly 300	Val	Gln	Leu	Lys
20	Lys 305	Gly	Asp	Gln	Ile	Leu 310	Leu	Pro	Gln	Met	Leu 315	Ser	Gly	Leu	Asp	Glu 320
	Arg	Glu	Asn	Ala	Cys 325	Pro	Met	His	Val	Asp 330	Phe	Ser	Arg	Gln	Lys 335	Val
25	Ser	His	Thr	Thr 340	Phe	Gly	His	Gly	Ser 345	His	Leu	Cys	Leu	Gly 350	Gln	His
30	Leu	Ala	Arg 355	Arg	Glu	Ile	Ile	Val 360	Thr	Leu	Lys	Glu	Trp 365	Leu	Thr	Arg
	Ile	Pro 370	Asp	Phe	Ser	Ile	Ala 375	Pro	Gly	Ala	Gln	Ile 380	Gln	His	Lys	Ser
35	Gly 385	Ile	Val	Ser	Gly	Val 390	Gln	Ala	Leu	Pro	Leu 395	Val	Trp	Asp	Pro	Ala 400
	Thr	Thr	Lys		Val 405						•					
40				The I	ONA	mole	cule	enco	dina	D 450	L.	oo th		1	J	quence of
	SEQ.	ID.					ouic	CHCO	ung	F430	cam II	as int	: nuc	ieotic	ie sec	quence of
45	ctgca	ggat	c gtt	atcc	gct g	gccg	atcto	atc	accc	agc g	rtttt	ccat	. cga	cgagg	gcc 6	0
	agcaa	ggca	c ttg	aact	ggt c	aagg	cagga	gca	ctga	tca a	accc	gtgat	cga	ctcca	act 1	20
ctttagccaa cccgcgttcc aggagaacaa caacaatgac gactgaaacc atacaaagca 180																

acgccaatct tgccctctg ccacccatg tgccagagca cctggtattc gacttcgaca 240
tgtacaatcc gtcgaatctg tctgccggcg tgcaggaggc ctgggcagtt ctgcaagaat 300
caaacgtacc ggatctggtg tggactcgct gcaacggcgg acactggatc gccactcgcg 360

gecaactgat cegtgaggee tatgaagatt acegecactt teccagegag tgecegttea 420 teeetegtga ageeggegaa geetacgact teatteecae etegatgat eegeeegage 480

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Natl. Acad. Sci. USA, 268:19681-19689 (1993); Kempf "Truncated Human P450 2D6: Expression in *Excherichia coli*, Ni²⁺-chelate Affinity Purification, and Characterization of Solibility and Aggregation," <u>Arch. Biochem. Biophys.</u>, 321:277-288 (1995), which are hereby incorporated by reference).

Mutations or variants of the above fusion protein are encompassed by the present invention.

Variants may be modified by, for example, the deletion or addition of amino acids that have minimal influence on the properties, secondary structure and hydropathic nature of the polypeptide. For example, a polypeptide may be conjugated to a signal (or leader) sequence at the N-terminal end of the protein which cotranslationally or post-translationally directs transfer of the protein. The polypeptide may also be conjugated to a linker or other sequence for ease of synthesis, purification, or identification of the polypeptide.

The DNA molecule encoding the cytochrome P450 polypeptide can be incorporated in cells using conventional recombinant DNA technology. Generally, this involves inserting the DNA molecule into an expression system to which the DNA molecule is heterologous (i.e. not normally present). The heterologous DNA molecule is inserted into the expression system or vector in proper sense orientation and correct reading frame. The vector contains the necessary elements for the transcription and translation of the inserted protein-coding sequences.

U.S. Patent No. 4,237,224 to Cohen and Boyer, which is hereby incorporated by reference, describes the production of expression systems in the form of recombinant plasmids using restriction enzyme cleavage and ligation with DNA ligase. These recombinant plasmids are then introduced by means of transformation and replicated in unicellular cultures including procaryotic organisms and eucaryotic cells grown in tissue culture.

Recombinant genes may also be introduced into viruses, such as vaccina virus. Recombinant viruses can be generated by transfection of plasmids into cells infected with virus.

Suitable vectors include, but are not limited to, the following viral vectors such as lambda vector system gtl1, gt WES.tB, Charon 4, and plasmid vectors such as pBR322, pBR325, pACYC177, pACYC184, pUC8, pUC9, pUC18, pUC19,

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Similarly, translation of mRNA in procaryotes depends upon the presence of the proper procaryotic signals which differ from those of eucaryotes. Efficient translation of mRNA in procaryotes requires a ribosome binding site called the Shine-Dalgarno ("SD") sequence on the mRNA. This sequence is a short nucleotide sequence of mRNA that is located before the start codon, usually AUG, which encodes the amino-terminal methionine of the protein. The SD sequences are complementary to the 3'-end of the 16S rRNA (ribosomal RNA) and probably promote binding of mRNA to ribosomes by duplexing with the rRNA to allow correct positioning of the ribosome. For a review on maximizing gene expression see Roberts and Lauer, Methods in Enzymology, 68:473 (1979), which is hereby incorporated by reference.

Promotors vary in their "strength" (i.e. their ability to promote transcription). For the purposes of expressing a cloned gene, it is desirable to use strong promotors in order to obtain a high level of transcription and, hence, expression of the gene. Depending upon the host cell system utilized, any one of a number of suitable promotors may be used. For instance, when cloning in *E. coli*, its bacteriophages, or plasmids, promotors such as the T7 phage promoter, *lac* promotor, *trp* promotor, *rec*A promotor, ribosomal RNA promotor, the P_R and P_L promotors of coliphage lambda and others, including but not limited, to *lac*UV5, *omp*F, *bla*, *lpp*, and the like, may be used to direct high levels of transcription of adjacent DNA segments. Additionally, a hybrid *trp-lac*UV5 (*tac*) promotor or other *E. coli* promotors produced by recombinant DNA or other synthetic DNA techniques may be used to provide for transcription of the inserted gene.

Bacterial host cell strains and expression vectors may be chosen which inhibit the action of the promotor unless specifically induced. In certain operons, the addition of specific inducers is necessary for efficient transcription of the inserted DNA. For example, the *lac* operon is induced by the addition of lactose or IPTG (isopropylthio-beta-D-galactoside). A variety of other operons, such as *trp*, *pro*, etc., are under different controls.

Specific initiation signals are also required for efficient gene transcription and translation in procaryotic cells. These transcription and translation initiation signals may vary in "strength" as measured by the quantity of gene specific

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mismatch will not hybridize (conditions of 85% stringency), and more preferably still, stringent conditions are those under which DNA sequences with more than 10% mismatch will not hybridize (conditions of 90% stringency). In a most preferred embodiment, stringent conditions are those under which DNA sequences with more than 6% mismatch will not hybridize (conditions of 94% stringency).

In yet another aspect of the present invention, the fusion protein can be applied to an environmental pollutant, such as an insecticide or other halogenated hydrocarbon spills, as part of a method of bioremediation. In fact, P450 enzymes can oxidize almost any compound that has a carbon-hydrogen bond and, thus, are useful for almost any environmental contaminant. Generally, microorganisms are extremely useful as agents for clean-up of environmental problems. Development of suitable microorganisms involves either selecting microorganisms with a bioremediation trait or by introducing a gene into microbes to engender them with that ability. By introducing the chimeric DNA molecule into an appropriate vector, it is possible to achieve bioremediation of environmental pollutants. Suitable vectors are non-pathogenic bacteria.

Another aspect of the present invention is using the fusion protein in a process of hydroxylating a compound to be oxidized. Typical compounds to be oxidized include hydrocarbons or any compound having a carbon-hydrogen bond. As discussed above, this involves contacting the compound to be oxidized with the fusion protein under conditions effective to hydroxylate the compound to be oxidized. The fusion protein can be provided by introducing the chimeric DNA molecule into an appropriate vector to express the fusion protein. Suitable vectors include pcW or pkk233-2.

Typicaly, hydroxylation occurs at from about 30 to about 50°C, with 37°C being preferred, with a potassium phosphate buffer and KCl (pH 7.4). The reaction can be monitored by the addition of dichloromethane and assaying by gas chromatography/mass spectrometry.

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EXAMPLES

The following examples illustrate, but are not intended to limit, the present invention.

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primer 7 CATCACCATCACCATCACTGAAGAAGAGCAGATGGCCTGGC (SEQ. ID. No. 13)
primer 8 GACAGGAATGAAGCACAGCTGGTA (SEQ. ID. No. 14)

5 Example 2 - Expression of the Fusion Protein

A single ampicillin-resistant colony of DH5α cells transformed with plasmid DNA was grown overnight at 37°C in Luria-Bertani medium containing 100 μg ampicillin ml⁻¹. A 0.5-ml aliquot was used to innoculate 50 ml of Terrific broth ("TB") and cultured for 10 h. This aliquot of 25 ml was used to innoculate 500 ml of TB media. Incubation at 37°C was continued for 19 h. The TB media was supplemented with ampicillin (100 μg ml⁻¹), 0.2% glucose, 100 μM δ-aminolevulinic acid, vitamins (100⁻¹w/w, Basal Medium Eagle Vitamin Solution, Gibco BRL, Grand Island, NY), and trace elements (2 mM MgSO₄.7H₂O, 0.1 mM CaCl₂, 1.0 μM FeSO₄, metal solution1, 50 μM H₃BO₄, 0.2 μM CoCl₂.6H₂O, 1 mM CuSO₄.5H₂O, 1 mM MnCl₂.4H₂O, 1 nM Na₂MoO₄ and 2 mM ZnCl₂). The cells were harvested by centrifugation at 5,000 g and 4°C for 10 min. The pellet was stored at -80°C before use.

20 Example 3 - Construction of Expression Plasmid for Pd and PdR

Nde I restriction site was introduced at the site of the initiation codon of the Pd or PdR plasmids by the procedures similar to those described above. After digestion of Pd by Sma I and digestion of PdR by Mlu I followed by blunt-ending, each plasmid was digested by Nde I. Gel purified DNA was cloned into PET-15, an expression vector (Novagene, Madison, WI), after digestion by Xho I and blunt-ending. E. coli strain BL21(DE3) was transformed with pETPd or pETPdR.

Pd and PdR were expressed as follows. Icoculum cultures (25 ml) of *E. coli* BL21(DE3), transformed with pETPd or pETPdR were grown at 37°C in M9 minimum medium supplemented with 100 µg ampicillin ml⁻¹, 0.5% glucose, vitamins, and trace elements as mentioned above. A 25-ml aliquot was used to inoculate 500 ml of M9 minimum medium and the flask was shaken for 1 h at 37°C, at which

Reductase and Puridaredoxin: Cloning, Sequence, and Heterologous Expression of the Proteins," J. Biol. Chem., 265:6066-6073 (1990), which is hereby incorporated by reference). Expression levels of the wild type P450_{cam} was 600-1000 nmoles/liter under similar conditions. After treatment with lysozyme and sonication of the cell pellet, the cell lysate was centrifuged at 105,000g and the supernatant was applied to a Ni-NTA agarose and hydroxylapatite columns (Imai et al., "Expression and Purification of Functional Human 17α -hydroxylase/17,20-lyase (P45017) in Escherichia coli," Proc. Natl. Acad. Sci. USA, 268:19681-19689 (1993), which is hereby incorporated by reference). The purified chimera showed a CO-reduced 10 difference spectrum at 448 nm (Fig. 2A) (Omura et al., "The Carbon Monoxide-Binding Pigment of Liver Microsomes I Evidence for its Hemeprotein Nature," J. Biol. Chem., 239:2370-2378 (1964), which is hereby incorporated by reference), and showed two major bands on SDS-polyacrylamide gel electrophoresis (Fig. 2B) (Laemmli, U.K., "Cleavage of Structural Protein During the Assembly of the Head of Bacteriophage," Nature, 227:680-685 (1970), which is hereby incorporated by 15 reference). Similar bands are observed from purified wild-type P450cam with a [His]6 tag coding sequence. The lower molecule weight band is presently unidentified. The resulting purified protein showed an approximae molecular weight of 51 kDa as judged by SDS-polyacrylamide gel electrophoresis, consistent with the 20 molecular weight expected for the chimera (Figure 2B).

The resulting pruified protein showed a reduced CO difference spectrum at 450 nm (Figure 2A). These data are consistent with a folded P450 protein having a functional active site. The observation that a functional chimera of P450 2C9 and P450_{cam}, which have only 15% primary sequence homology, can still bind CO provides strong evidence for a conserved three-dimensional structure between P450_{cam} and CYP2 family. The fact that the resulting enzyme is soluble, while mammalian enzymes with the amino terminus removed are not, indicates that other regions near the amino terminus may also be important for membrane interactions. (Lemos-Chiarandine et al., <u>J. Cell Biol.</u>, 104:209-219 (1987); Vergeres et al.,

Biochemistry, 28:3650-3655 (1989); Wachenfeldt et al., <u>Arch. Biochem. Biophys.</u>,

339:107-114 (1997), which are hereby incorporated by reference.)

putidaredoxin reductase, and 300 μ M NADH. The reaction was stopped by the addition of 4 ml of dichloromethane and assayed by gas chromatography/mass spectrometry. Experiments to determine if the mammalian P450 reductase can support the same oxidation are underway.

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Detection of the catalyic activity toward 4-chlorotoluene indicate that the fusion protein can function as an active P450 enzyme (Table 1). As compared with the turnover number from the wild type P450_{cam}, the chimera shows approximately 3 times the activity towards 4-chlorotoluene. This means a potential for making soluble P450 that can perform stereospecific synthesis.

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This approach could have a number of applications. 1) From other homology models of mammalian P450 enzymes it is apparent that this method may prove to be a general method for constructed soluble P450 enzymes with mammalian active site characteristics. These enzymes should be more adaptable to uses in benign synthesis and bioremediation than the more restrictive bacterial enzymes and easier to work with then the membrane bound mammalian enzymes. 2) Selectively replacing amino acid segments in the amino terminus with the mammalian amino acids may prove to be a valuable method of determining important membrane association sites.

3) Since the enzyme is soluble, it could prove a method for obtaining structural information. In particular it should be amiable to Xray crystallography. 4) Since the enzyme is part mammalian and part bacterial, it can be used to determine the features that confer specific interactions with the different reductases system that are used by the bacterial and mammalian proteins.

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Although the invention has been described in detail for the purpose of illustration, it is understood that such detail is solely for that purpose, and variations can be made therein by those skilled in the art without departing from the spirit and scope of the invention which is defined by the following claims.

- 8. A chimeric DNA molecule according to claim 1, wherein the chimeric DNA molecule has a heme ligand positioned in a relative orientation to an I-helix and a fifth cysteine ligand similar to that of the heme ligand in a full length mammalian P450 protein.
- 9. A chimeric DNA molecule according to claim 1, wherein the chimeric DNA molecule encodes an amino acid sequence of SEQ. ID. No. 2.
- 10. A chimeric DNA molecule according to claim 9, wherein the chimeric DNA molecule has a nucleotide sequence of SEQ. ID. No. 1.
 - 11. A DNA expression system transformed with the chimeric DNA molecule of claim 1.
 - 12. A DNA expression system according to claim 11, wherein the chimeric DNA molecule is positioned in the expression system in proper sense orientation and correct reading frame.
- 20 13. A DNA expression system according to claim 11, wherein the first and second DNA molecules are fused together at a location where the encoded fusion protein lacks secondary structure.
- 14. A host cell transformed with the chimeric DNA molecule of claim 1.
 - 15. A host cell according to claim 14, wherein the host cell is selected from the group consisting of plant cells, mammalian cells, insect cells, and bacterial cells.
- 16. A fusion protein comprising:

 a portion of a bacterial P450 protein and
 a portion of a mammalian P450 protein fused to the portion of a bacterial P450 protein, wherein the fusion protein is active and soluble in aqueous

 35 liquid.

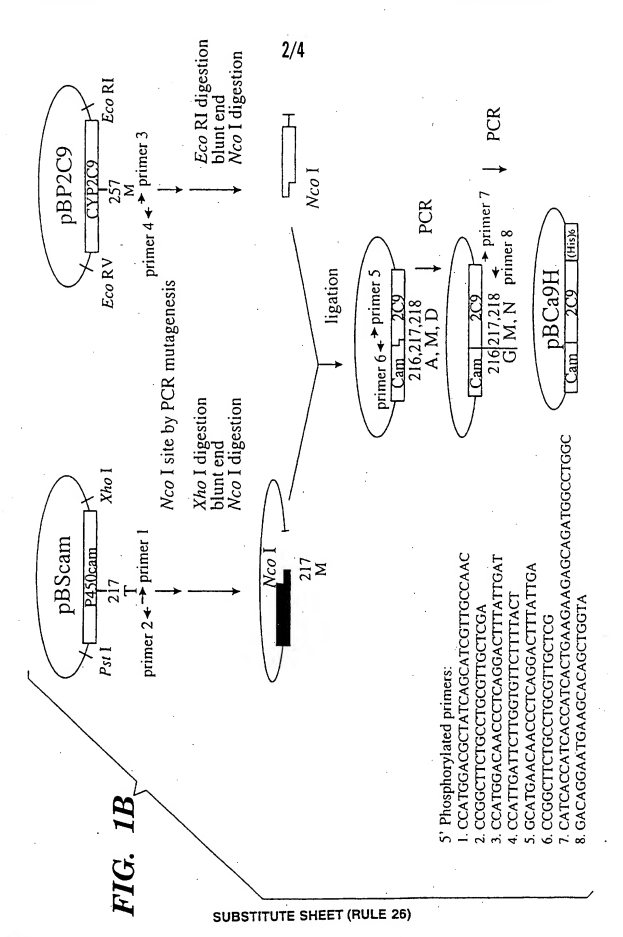
contacting the compound to be oxidized with the fusion protein according to claim 16 under conditions effective to hydroxylate the compound to be oxidized.

- 5 26. A method according to claim 25, wherein the portion of the mammalian P450 protein and the portion of the bacterial P450 protein are fused where the encoded fusion protein lacks secondary structure.
- 27. A method according to claim 25, wherein the fusion protein is prepared from a full length mammalian P450 protein where a portion of the full length mammalian P450 protein is replaced with a homologous portion of a full length bacterial P450 protein.
- 28. A method according to claim 27, wherein all amino acids prior to a random coil between G- and H-helices in the full length mammalian P450 protein are replaced with a homologous portion of the full length bacterial P450 protein.
- 29. A method according to claim 27, wherein the fusion protein comprises about 50 percent of the full length mammalian P450 protein and about 50 percent of the full length bacterial P450 protein.
 - 30. A method according to claim 25, wherein the fusion protein is provided by providing a vector comprising a chimeric DNA molecule comprising:

 a first DNA molecule encoding a portion of a full length
- 25 bacterial P450 protein;
 - a second DNA molecule fused to the first DNA molecule and encoding a portion of a full length mammalian P450 protein, wherein the chimeric DNA molecule encodes the fusion protein.
- 31. A method according to claim 30, wherein the first and second DNA molecules are fused together at a location where the encoded fusion protein lacks secondary structure.
- 32. A method according to claim 30, wherein the chimeric DNA molecule is prepared from a DNA molecule encoding a full length mammalian P450

- 40. A method according to claim 35, wherein the fusion protein is provided by providing a vector comprising a chimeric DNA molecule comprising:

 a first DNA molecule encoding a portion of a full length bacterial P450 protein;
- a second DNA molecule fused to the first DNA molecule and encoding a portion of a full length mammalian P450 protein, wherein the chimeric DNA molecule encodes the fusion protein.
- 41. A method according to claim 40, wherein the first and second
 DNA molecules are fused together at a location where the encoded fusion protein lacks secondary structure.
- 42. A method according to claim 40, wherein the chimeric DNA molecule is prepared from a DNA molecule encoding a full length mammalian P450
 protein where a portion of the DNA molecule encoding a full length mammalian P450 protein is replaced with a DNA molecule encoding a homologous portion of a full length bacterial P450 protein.
- 43. A method according to claim 42, wherein all amino acids prior to a random coil between G- and H-helices in the full length mammalian P450 protein are replaced with a homologous portion of the full length bacterial P450 protein.
- 44. A method according to claim 42, wherein the chimeric DNA molecule comprises about 50 percent of the DNA molecule encoding the full length
 25 mammalian P450 protein and about 50 percent of the DNA molecule encoding the full length bacterial P450 protein.



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4/4

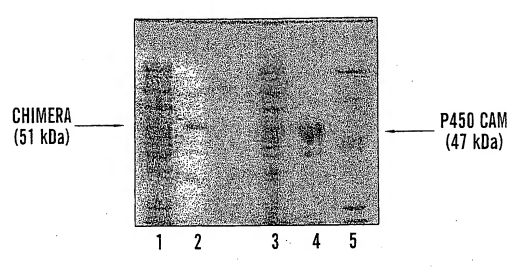


FIG. 2B

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Trp Ala Val Leu Gln Glu Ser Asn Val Pro Asp Leu Val Trp Thr Arg
35 40 45

Cys Asn Gly Gly His Trp Ile Ala Thr Arg Gly Gln Leu Ile Arg Glu
50 55 60

Ala Tyr Glu Asp Tyr Arg His Phe Ser Ser Glu Cys Pro Phe Ile Pro 65. 70 75 80

Arg Glu Ala Gly Glu Ala Tyr Asp Phe Ile Pro Thr Ser Met Asp Pro 85 90 95

Pro Glu Gln Arg Gln Phe Arg Ala Leu Ala Asn Gln Val Val Gly Met 100 \$105\$

Pro Val Val Asp Lys Leu Glu Asn Arg Ile Gln Glu Leu Ala Cys Ser 115 120 125

Leu Ile Glu Ser Leu Arg Pro Gln Gly Gln Cys Asn Phe Thr Glu Asp 130 135 140

Pro Glu Glu Asp Ile Pro His Leu Lys Tyr Leu Thr Asp Gln Met Thr
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Arg Pro Asp Gly Ser Met Thr Phe Ala Glu Ala Lys Glu Ala Leu Tyr 180 185 190 WO 99/08812 PCT/US98/16979

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Pro Thr Pro Leu Pro Val Ile Gly Asn Ile Leu Gln Ile Gly Ile Lys
35 40 45

Asp Ile Ser Lys Ser Leu Thr Asn Leu Ser Lys Val Tyr Gly Pro Val 50 60

Phe Thr Leu Tyr Phe Gly Leu Lys Pro Ile Val Val Leu His Gly Tyr 65 70 75 80

Glu Ala Val Lys Glu Ala Leu Ile Asp Leu Gly Glu Glu Phe Ser Gly 85 90 95

Arg Gly Ile Phe Pro Leu Ala Glu Arg Ala Asn Arg Gly Phe Gly Ile 100 105 110

Val Phe Ser Asn Gly Lys Lys Trp Lys Glu Ile Arg Arg Phe Ser Leu 115 120 125

Met Thr Leu Arg Asn Phe Gly Met Gly Lys Arg Ser Ile Glu Asp Arg 130 135 140

Ala Ser Pro Cys Asp Pro Thr Phe Ile Leu Gly Cys Ala Pro Cys Asn 165 170 175.

Val Ile Cys Ser Ile Ile Phe His Lys Arg Phe Asp Tyr Lys Asp Gln 180 185 190

Gln Phe Leu Asn Leu Met Glu Lys Leu Asn Glu Asn Ile Lys Ile Leu 195 200 205

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Lys Gly Asp Gln Ile Leu Leu Pro Gln Met Leu Ser Gly Leu Asp Glu 305 310 315 320

Arg Glu Asn Ala Cys Pro Met His Val Asp Phe Ser Arg Gln Lys Val 325 330 335

Ser His Thr Thr Phe Gly His Gly Ser His Leu Cys Leu Gly Gln His 340 345 350

Leu Ala Arg Arg Glu Ile Ile Val Thr Leu Lys Glu Trp Leu Thr Arg 355 360 . 365

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16979

A. CLASSIFICATION OF SUBJECT MATTER IPC(6): B09B 3/00; C12N 1/00, 5/10, 9/02, 15/53; 15/63; C12P 1/00, 7/02 US CL: Please See Extra Sheet.							
	o International Patent Classification (IPC) or to both	national classification and IPC					
	DS SEARCHED						
	ocumentation searched (classification system follower						
U.S. :	536/23.2; 23.7; 435/41, 56, 57, 58, 59, 61,125, 189,	, 262.5, 69.1, 320.1, 252.3, 254.11, 325,	410				
Documentati	ion searched other than minimum documentation to th	e extent that such documents are included	in the fields searched				
Electronic d	ata base consulted during the international search (n	ame of data base and, where practicable,	search terms used)				
Please Sec	Extra Sheet.		·				
C. DOC	UMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where as	ppropriate, of the relevant passages	Relevant to claim No.				
A	US 5,114,852 A (YABUSAKI et document.	al.) 19 May 1992, entire	1-44				
A	US 5,240,831 A, (H.J. BARNES) 31 August 1993, entire document. 1-44						
A	O'KEEFE et al. Occurrence and biological function of cytochrome P450 monooxygenases in the actinomycetes. Molecular Microbiology. 1991. Vol. 5, No. 9, pages 2099-2105, entire document.						
A	OKUDA et al. Recent progress in biology of enzymes involved in vitam Lipid Research. 1995. Vol. 36, pages	in D metabolism. Journal of	25-34				
X Furth	er documents are listed in the continuation of Box C	See patent family annex.					
• Spe	soial categories of cited documents:	"I" later document published after the inte- date and not in conflict with the appli	rnational filing date or priority ication but cited to understand				
	nument defining the general state of the art which is not considered be of particular relevance	the principle or theory underlying the	invention				
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cita	nament which may throw doubts on priority claim(s) or which is do establish the publication date of another citation or other cial reason (as specified)	when the document is taken alone 4Ye document of perticular relevance; the	claimed invention cannot be				
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18 SEPTE	MBER 1998	20001	770				
Commission Box-PCT	Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Authorized officer GABRIELE E. BUGAISKY						
Facsimile No		Telephone No. (703) 308-0196	/				

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US98/16979

A. CLASSIFICATION OF SUBJECT US CL:	MATTER:
536/23.2; 435/41, 189, 262.5, 69.1, 32	20.1, 252.3, 254.11, 325, 410
B. FIELDS SEARCHED Electronic data bases consulted (Name	of data base and where practicable terms used):
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